# PCT

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I) Applicant: ORTHO PHARMACEUTICAL CORPO [US/US]; U.S. Route #202, P.O. Box 300, Raritan, I (US).	RATION NJ 08869	Published  Without international search requestion receipt of that report.	port and to be republished			
72) Inventors: SHANK, Richard, P.; 551 Village Circle, E PA 19422-1636 (US). DERIAN, Claudia, K.; 104 I Road, Hatboro, PA 19040 (US).	Blue Bell, East Mill		j			
74) Agents: CIAMPORCERO, Audiey, A., Jr. et al.; Johnson, One Johnson & Johnson Plaza, New Bri NJ 08933-7003 (US).	hnson & unswick,					
4) Title: ANTICONVULSANT DERIVATIVES USEFU. 7) Abstract	L IN TR	EATING PSORIASIS				
Anticonvulsant derivatives useful in treating psoriasis	are discl	osed.				
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#### ANTICONVULSANT DERIVATIVES USEFUL IN TREATING PSORIASIS

#### **BACKGROUND OF THE INVENTION**

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### Compounds of Formula I:

$$R_5$$
 $R_4$ 
 $R_2$ 
 $R_3$ 
 $R_2$ 

10 are structurally novel antiepileptic compounds that are highly effective anticonvulsants in animal tests (Maryanoff, B.E., Nortey, S.O., Gardocki, J.F., Shank, R.P. and Dodgson, S.P. J. Med. Chem. 30, 880-887, 1987; Maryanoff. B.E., Costanzo, M.J., Shank, R.P., Schupsky, J.J., Ortegon, M.E., and Vaught J.L. Bioorganic & Medicinal Chemistry Letters 3, 2653-2656, 1993, McComsey, 15 D.F. and Maryanoff, B.E., J. Org. Chem. 1995). These compounds are covered by US Patent No. 4,513,006. One of these compounds 2,3:4,5-bis-O-(1methylethylidene)-B-D-fructopyranose sulfamate known as topiramate has been demonstrated in clinical trials of human epilepsy to be effective as adjunctive therapy or as monotherapy in treating simple and complex partial seizures and 20 secondarily generalized seizures (E. FAUGHT, B.J. WILDER, R.E. RAMSEY, R.A. REIFE, L.D. KRAMER, G.W. PLEDGER, R.M. KARIM et. al., Epilepsia 36 (S4) 33, 1995; S.K. SACHDEO, R.C. SACHDEO, R.A. REIFE, P. LIM and G. PLEDGER, Epilepsia 36 (S4) 33, 1995), and is currently marketed for the treatment of simple and complex partial seizure epilepsy with or without 25 secondary generalized seizures in Great Britain, Finland, the United States and Sweden and applications for regulatory approval are presently pending in numerous countries throughout the world.

Compounds of Formula I were initially found to possess anticonvulsant activity in the traditional maximal electroshock seizure (MES) test in mice

(SHANK, R.P., GARDOCKI, J.F., VAUGHT, J.L., DAVIS, C.B., SCHUPSKY, J.J., RAFFA, R.B., DODGSON, S.J., NORTEY, S.O., and MARYANOFF, B.E., Epilepsia 35 450-460, 1994). Subsequent studies revealed that Compounds of Formula I were also highly effective in the MES test in rats. More recently topiramate was found to effectively block seizures in several rodent models of epilepsy (J. NAKAMURA, S. TAMURA, T. KANDA, A. ISHII, K. ISHIHARA, T. SERIKAWA, J. YAMADA, and M. SASA, Eur. J. Pharmacol. 254 83-89, 1994), and in an animal model of kindled epilepsy (A. WAUQUIER and S. ZHOU, Epilepsy Res. 24, 73-77, 1996).

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Recent preclinical studies on topiramate have revealed previously unrecognized pharmacological properties which suggest that topiramate should be effective in treating some other disorders. One of these is psoriasis.

### 15 DISCLOSURE OF THE INVENTION

Accordingly, it has been found that compounds of the following formula I:

$$R_5$$
 $R_4$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_3$ 

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wherein X is O or CH<sub>2</sub>, and R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub> and R<sub>5</sub> are as defined hereinafter are useful in treating psoriasis.

### 25 DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The sulfamates of the invention are of the following formula (I):

$$R_5$$
 $R_4$ 
 $R_3$ 
 $R_2$ 
 $R_3$ 
 $R_2$ 

wherein

5 X is CH<sub>2</sub> or oxygen;

R<sub>1</sub> is hydrogen or alkyl; and

R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub> and R<sub>5</sub> are independently hydrogen or lower alkoxy, when X is oxygen, R<sub>2</sub> and R<sub>3</sub> and/or R<sub>4</sub> and R<sub>5</sub> together may be a methylenedioxy group of the following formula (II):

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wherein

R6 and R7 are the same or different and are hydrogen, lower alkyl or are alkyl and are joined to form a cyclopentyl or cyclohexyl ring.

R<sub>1</sub> in particular is hydrogen or alkyl of about 1 to 4 carbons, such as methyl, ethyl and iso-propyl. Alkyl throughout this specification includes straight and branched chain alkyl. Alkyl groups for R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub>, R<sub>5</sub>, R<sub>6</sub> and R<sub>7</sub> are of about 1 to 3 carbons and include methyl, ethyl, iso-propyl and n-propyl.

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A particular group of compounds of formula (I) are those wherein X is oxygen and both R2 and R3, and R4 and R5 together are methylenedioxy groups of the formula (II), wherein R6 and R7 are both hydrogen, both alkyl, or combine to form a spiro cyclopentyl or cyclohexyl ring, in particular where R6 and R7 are both alkyl such as methyl. A second group of compounds are those wherein X is CH2 and R4 and R5 are joined to form a benzene ring. A third

group of compounds of formula (I) are those wherein both R2 and R3 are hydrogen.

The compounds of formula (I) may be synthesized by the following methods:

(a) Reaction of an alcohol of the formula RCH2OH with a chlorosulfamate of the formula CISO2NH2 or CISO2NHR1 in the presence of a base such as potassium a-butoxide or sodium hydride at a temperature of about -20° to 25° C and in a solvent such as toluene, THF or dimethylformamide wherein R is a moiety of the following formula (III):

$$R_5$$
 $R_4$ 
 $R_2$ 
 $R_3$ 

15 (b) Reaction of an alcohol of the formula RCH2OH with sulfurylchloride of the formula SO2Cl2 in the presence of a base such as triethylamine or pyridine at a temperature of about -40° to 25° C in a solvent such as diethyl ether or methylene chloride to produce a chlorosulfate of the formula RCH2OSO2Cl.

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The chlorosulfate of the formula RCH<sub>2</sub>OSO<sub>2</sub>Cl may then be reacted with an amine of the formula R<sub>1</sub>NH<sub>2</sub> at a temperature of abut 40° to 25° C in a solvent such as methylene chloride or acetonitrile to produce a compound of formula (I). The reaction conditions for (b) are also described by T. Tsuchiya et al. in Tet. Letters, No. 36, p. 3365 to 3368 (1978).

(c) Reaction of the chlorosulfate RCH<sub>2</sub>OSO<sub>2</sub>Cl with a metal azide such as sodium azide in a solvent such as methylene chloride or acetonitrile yields an azidosulfate of the formula RCH<sub>2</sub>OSO<sub>2</sub>N<sub>3</sub> as described by M.

Hedayatullah in Tet. Lett. p. 2455-2458 (1975). The azidosulfate is then reduced to a compound of formula (I) wherein R<sub>1</sub> is hydrogen by catalytic hydrogenation, e.g. with a noble metal and H<sub>2</sub> or by heating with copper metal in a solvent such as methanol.

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The starting materials of the formula RCH2OH may be obtained commercially or as known in the art. For example, starting materials of the formula RCH2OH wherein both R2 and R3, and R4 and R5 are identical and are of the formula (II) may be obtained by the method of R. F. Brady in Carbohydrate Research, Vol. 14, p. 35 to 40 (1970) or by reaction of the trimethylsilyl enol ether of a R6COR7 ketone or aldehyde with fructose at a temperature of about 25° C, in a solvent such a halocarbon, e.g. methylene chloride in the presence of a protic acid such as hydrochloric acid or a Lewis Acid such as zinc chloride. The trimethylsilyl enol ether reaction is described by G. L. Larson et al in J. Org. Chem. Vol. 38, No. 22, p. 3935 (1973).

Further, carboxylic acids and aldehydes of the formulae RCOOH and RCHO may be reduced to compounds of the formula RCH2OH by standard reduction techniques, e.g. reaction with lithium aluminum hydride, sodium borohydride or borane-THF complex in an inert solvent such a diglyme, THF or toluene at a temperature of about 0° to 100° C, e.g. as described by H.O. House in "Modern Synthetic Reactions", 2nd Ed., pages 45 to 144 (1972).

The compounds of formula I: may also be made by the process disclosed US Patent: No.4,513,006, which is incorporated by reference herein.

The compounds of formula I include the various individual isomers as well as the racemates thereof, e.g., the various alpha and beta attachments, i.e., below and above the plane of the drawing, of R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub> and R<sub>5</sub> on the 6-membered ring. Preferably, the oxygens of the methylenedioxy group (II) are attached on the same side of the 6-membered ring.

The activity of the compounds of formula I in treating psoriasis was first evidenced in clinical studies conducted to evaluate the efficacy of topiramate in treating epilepsy. At least three patients who coincidentally had psoriasis reported that there was a marked reduction in the psoriatic lesions. Therefore, preclinical in vitro studies were conducted to evaluate the effects of topiramate on keratinocyte function as a putative mechanism of action for its potential beneficial effects in treating psoriasis. One of the hallmarks of psoriatic lesions is hyperproliferative epidermal keratinocytes. In general, agents that affect the prollferation of keratinocytes have an inverse effect on differentiation, i.e. they would inhibit growth and enhance differentiation. Two measures of keratinocyte function were therefore evaluated: cell growth and differentiation.

In these studies, keratinocytes were grown in Medium154, a low calcium medium supplemented with bovine pituitary extract (BPE), bovine insulin, bovine transferrin, human epidermal growth factor (EGF) and hydrocortisone. Keratinocytes were grown to 60-80% confluence and subcultured by using trypsin/EDTA.

Four separate experiments were performed to evaluate the dosedependent effect of topiramate on keratinocyte cell growth as measured by maturity after six days of treatment. Topiramate was dissolved in DMSO to make a 100 mM stock solution. In all experiments the final concentration of DMSO in the cell incubation medium was 0.1%. Vehicle controls (0.1% DMSO) were included in each experiment. Cell growth was induced by a combination of the growth factors EGF and BPE. Topiramate had a modest inhibitory effect on cell growth under these assay conditions; however, no dose dependence was observed (R.W Johnson Pharmaceutical Research Institute Laboratory Notebook No. 12183 and 12540). The maximal response was observed at 10 micromolar, 32 ± 10% inhibition. While there was a trend toward inhibition of cell growth, this did not reach statistical significance (p>0.05).

The effect of topiramate on keratinocyte differentiation was measured by the expression of transglutaminase-1 protein after three days of treatment. Three separate experiments were performed. Differentiation was evaluated in both low calcium and high calcium incubation conditions. An increase in differentiation would be most readily observed under low calcium conditions whereas an inhibition of differentiation could be detected under conditions of high calcium-induced differentiation. Topiramate caused a modest increase in transglutaminase-1 protein with both conditions, indicating an enhancing effect. A follow-up study was conducted which extended the incubation time to 5 days to look for further enhancement. No additional increases in transglutaminase-1 were observed in this latter study.

The results of these studies indicate that topiramate's effects on keratinocyte function are consistent with those expected for an agent that would affect the hyperproliferative keratinocyte response associated with psoriasis; inhibition of cell growth and enhancement of differentiation.

For treating psoriasis, a compound of formula (I) may be employed at a daily dosage in the range of about 50 to 400 mg administered orally, usually in two divided doses, for an average adult human. A unit dose would contain about 25 to 200 mg of the active ingredient. Alternatively, a compound of formula (I) may be administered topically to the affected area of the skin once or twice daily at a dosage in the range of 5 to 50 mg.

To prepare the pharmaceutical compositions of this invention, one or more sulfamate compounds of formula (I) are intimately admixed with a pharmaceutical carrier according to conventional pharmaceutical compounding techniques, which carrier may take a wide variety of forms depending on the form of preparation desired for administration, e.g., oral, by suppository, or parenteral. In preparing the compositions in oral dosage form, any of the usual pharmaceutical media may be employed. Thus, for liquid oral preparations, such as for example, suspensions, elixirs and solutions, suitable carriers and

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additives include water, glycols, oils, alcohols, flav ring agents, preservatives, coloring agents and the like; for solid oral preparations such as, for example, powders, capsules and tablets, suitable carriers and additives include starches, sugars, diluents, granulating agents, lubricants, binders, disintegrating agents and the like. Because of their ease in administration, tablets and capsules represent the most advantageous oral dosage unit form, in which case solid pharmaceutical carriers are obviously employed. If desired, tablets may be sugar coated or enteric coated by standard techniques. Suppositories may be prepared, in which case cocoa butter could be used as the carrier. For parenterals, the carrier will usually comprise sterile water, though other ingredients, for example, for purposes such as aiding solubility or for preservation, may be included. Injectable suspensions may also be prepared in which case appropriate liquid carriers, suspending agents and the like may be employed.

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Topiramate is currently available for oral administration in round tablets containing 25 mg, 100 mg or 200 mg of active agent. The tablets contain the following inactive ingredients: lactose hydrous, pregelatinized starch, microcrystalline cellulose, sodium starch glycolate, magnesium stearate, purified water, carnauba wax, hydroxypropyl methylcellulose, titanium dioxide, polyethylene glycol, synthetic iron oxide, and polysorbate 80.

The pharmaceutical compositions herein will contain, per dosage unit, e.g., tablet, capsule, powder injection, teaspoonful, suppository and the like from about 25 to about 200 mg of the active ingredient.

#### WHAT IS CLAIMED IS:

1. A method for treating psoriasis comprising administering to a human afflicted with such condition a therapeutically effective amount for treating such condition of a compound of the formula I:

$$R_5$$
 $R_4$ 
 $R_2$ 
 $R_3$ 
 $R_2$ 

wherein

5 X is CH<sub>2</sub> or oxygen;

R<sub>1</sub> is hydrogen or alkyl; and

R2, R3, R4 and R5 are independently hydrogen or lower alkyl and, whin X is CH2, R4 and R5 may be alkene groups joined to form a benzene ring and, when X is oxygen, R2 and R3 and/or R4 and R5 together may be a methylenedioxy group of the following formula (II):

wherein

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15 R6 and R7 are the same or different and are hydrogen, lower alkyl or are alkyl and are joined to form a cyclopentyl or cyclohexyl ring.

- 2. The method of claim 1 wherein the compound of formula I is topiramate.
- The method of claim 1, wherein the therapeutically effective amount is of from about 50 to 400 mg.
  - 4. The method of claim 1, wherein the amount is of from about 25 to 200 mg.

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### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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(21) International Application Number: PCT/US (22) International Filing Date: 24 June 1997 (27) (30) Priority Data: 60/022,005 28 June 1996 (28.06.96) (71) Applicant: ORTHO PHARMACEUTICAL CORPOLIUS/US); U.S. Route #202, P.O. Box 300, Raritan, I (US). (72) Inventors: SHANK, Richard, P.; 551 Village Circle, E PA 19422-1636 (US). DERIAN, Claudia, K.; 104 Road, Hatboro, PA 19040 (US). (74) Agents: CIAMPORCERO, Audley, A., Jr. et al.; Jo Johnson, One Johnson & Johnson Plaza, New Br NJ 08933-7003 (US).	24.06.9  RATIO NJ 0886 Blue Be East M	BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA UG, UZ, VN, European patent (AT, BE, CH, DE, DK, ES FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).  Published With international search report. Before the expiration of the time limit for amending the claim and to be republished in the event of the receipt of amendments  (88) Date of publication of the international search report: 12 February 1998 (12.02.98)

(57) Abstract

Anticonvulsant derivatives useful in treating psoriasis are disclosed.

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# INTERNATIONAL SEARCH REPORT

International Application No. 97/10891

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C. DOCUM	MENTS CONSIDERED TO BE RELEVANT							
Category *	Citation of document, with indication, where appropriate, of the r	elevant passages	Relevant to claim No.					
A	US 4513006 A		1-4					
**	(MARYANOFF, B.E. et a	11.)						
	23 April 1985 (23.04.	.85),						
	abstract, claims 5-9,							
	column 1, lines 16-33 column 3, line 15 - 6	rolumn 5.						
	line 19, example 3							
	(cited in the application)	ation).						
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A	US 4792569 A	-2 \	1,3,4					
ļ	(MARYANOFF, B.E. et a 20 December 1988 (20	12.88).						
	abstract, claims 1,10	0-12,						
	column 1, lines 20-3	7,						
	column 3, line 29 - c	column 4,						
	line 33.							
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# INTERNATIONAL SEARCH REPORT

inter onal application No.

PCT/US 97/10891

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Box I Obs rvati ns where ertain claims were f und unsearchab	
This International Search Report has not been established in respect of certain o	laims under Article 17(2)(a) for the following reasons:
1. X Claims Nos.: because they relate to subject matter not required to be searched by the Remark: Although claim(s) 1-4 is(are) directed to a method of to body, the search has been carried effects of the compound/composition  2. Claims Nos.: because they relate to parts of the international Application that do not an extent that no meaningful international Search can be carried out, a	reatment of the human/animal out and based on the alleged on.
Claims Nos.:     because they are dependent claims and are not drafted in accordance.	
Box II Observations where unity of invention is lacking (Continu	estion of item 2 of first sneet)
This international Searching Authority found multiple inventions in this internat	ional application, as follows:
As all required additional search fees were timely paid by the application of the searchable claims.	ant, this international Search Report covers all
As all searchable claims could be searched without effort justifying of any additional fee.	an additional fee, this Authority did not invite payment
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4. No required additional search fees were timely paid by the applica restricted to the invention first mentioned in the claims; it is covere	nt. Consequently, this International Search Report is d by claims Nos.:
Remark on Protest	al search fees were accompanied by the applicant's protest.

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#### ANHANG

### ANNEX

#### ANNEXE

zum internationalen Recherchen-bericht über die internationale Patentanmeldung Nr.

to the International Search Report to the International Patent Application No.

au rapport de recherche inter-national relatif à la demande de brevet international n°

### PCT/US 97/10891 SAE 168598

In diesem Anhang sind die Mitglieder der Patentfamilien der im oberigen nannten internationalen Recherchenbericht angeführten Patentdokumente angegeben. Diese Angaben dienen nur zur Linternichtung und erfolgen ohne Gewähr.

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# INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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(21) International Application Number: PCT/US  (22) International Filing Date: 24 June 1997 (  (30) Priority Data: 60/022,005 28 June 1996 (28.06.96)  (71) Applicant: ORTHO PHARMACEUTICAL CORPO [US/US]; U.S. Route #202, P.O. Box 300, Raritan, (US).  (72) Inventors: SHANK, Richard, P.; 551 Village Circle, FPA-19422-1636 (US). DERIAN, Claudia, K.; 104 Road, Hatboro, PA 19040 (US).  (74) Agents: CIAMPORCERO, Audley, A., Jr. et al.; Jo Johnson, One Johnson & Johnson Plaza, New Br NJ 08933-7003 (US).	(24.06.9 RATIO NJ 0886 Blue Bel East Mi shnson & unswick	BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).  Published  With international search report.  Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.  [1]  (88) Date of publication of the international search report:  12 February 1998 (12.02.98)
54) Title: ANTICONVULSANT DERIVATIVES USEFU	LINT	REATING PSORIASIS

### (57) Abstract

Anticonvulsant derivatives useful in treating psoriasis are disclosed.

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# ANTICONVULSANT DERIVATIVES USEFUL IN TREATING PSORIASIS

### BACKGROUND OF THE INVENTION

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### Compounds of Formula I:

$$R_5$$
 $R_4$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_3$ 
 $R_4$ 
 $R_3$ 

are structurally novel antiepileptic compounds that are highly effective 10 anticonvulsants in animal tests (Maryanoff, B.E, Nortey, S.O., Gardocki, J.F., Shank, R.P. and Dodgson, S.P. J. Med. Chem. 30, 880-887, 1987; Maryanoff, B.E., Costanzo, M.J., Shank, R.P., Schupsky, J.J., Ortegon, M.E., and Vaught J.L. Bioorganic & Medicinal Chemistry Letters 3, 2653-2656, 1993, McComsey, D.F. and Maryanoff, B.E., J. Org. Chem. 1995). These compounds are covered 15 by US Patent No. 4,513,006. One of these compounds 2,3:4,5-bis-O-(1methylethylidene)-B-D-fructopyranose sulfamate known as topiramate has been demonstrated in clinical trials of human epilepsy to be effective as adjunctive therapy or as monotherapy in treating simple and complex partial seizures and secondarily generalized seizures (E. FAUGHT, B.J. WILDER, R.E. RAMSEY, 20 R.A. REIFE, L D. KRAMER, G.W. PLEDGER, R.M. KARIM et. al., Epilepsia 36 (S4) 33, 1995; S.K. SACHDEO, R.C. SACHDEO, R.A. REIFE, P. LIM and G. PLEDGER, Epilepsia 36 (S4) 33, 1995), and is currently marketed for the treatment of simple and complex partial seizure epilepsy with or without secondary generalized seizures in Great Britain, Finland, the United States and 25 Sweden and applications for regulatory approval are presently pending in numerous countries throughout the world.

Compounds of Formula I were initially found to possess anticonvulsant activity in the traditional maximal electroshock seizure (MES) test in mice

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(SHANK, R.P., GARDOCKI, J.F., VAUGHT, J.L., DAVIS, C.B., SCHUPSKY, J.J., RAFFA, R.B., DODGSON, S.J., NORTEY, S.O., and MARYANOFF, B.E., Epilepsia 35 450-460, 1994). Subsequent studies revealed that Compounds of Formula I were also highly effective in the MES test in rats. More recently topiramate was found to effectively block seizures in several rodent models of epilepsy (J. NAKAMURA, S. TAMURA, T. KANDA, A. ISHII, K. ISHIHARA, T. SERIKAWA, J. YAMADA, and M. SASA, Eur. J. Pharmacol. 254 83-89, 1994), and in an animal model of kindled epilepsy (A. WAUQUIER and S. ZHOU, Epilepsy Res. 24, 73-77, 1996).

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Recent preclinical studies on topiramate have revealed previously unrecognized pharmacological properties which suggest that topiramate should be effective in treating some other disorders. One of these is psoriasis.

### 15 DISCLOSURE OF THE INVENTION

Accordingly, it has been found that compounds of the following formula I:

$$R_5$$
 $R_4$ 
 $R_3$ 
 $R_2$ 
 $R_3$ 

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wherein X is O or CH<sub>2</sub>, and R<sub>1</sub>, R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub> and R<sub>5</sub> are as defined hereinafter are useful in treating psoriasis.

### 25 DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The sulfamates of the invention are of the following formula (I):

$$R_5$$
 $R_4$ 
 $R_2$ 
 $R_3$ 
 $R_2$ 

wherein

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X is CH2 or oxygen;

R<sub>1</sub> is hydrogen or alkyl; and

R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub> and R<sub>5</sub> are independently hydrogen or lower alkoxy, when X is oxygen, R<sub>2</sub> and R<sub>3</sub> and/or R<sub>4</sub> and R<sub>5</sub> together may be a methylenedioxy group of the following formula (II):

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wherein

R6 and R7 are the same or different and are hydrogen, lower alkyl or are alkyl and are joined to form a cyclopentyl or cyclohexyl ring.

R<sub>1</sub> in particular is hydrogen or alkyl of about 1 to 4 carbons, such as methyl, ethyl and iso-propyl. Alkyl throughout this specification includes straight and branched chain alkyl. Alkyl groups for R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub>, R<sub>5</sub>, R<sub>6</sub> and R<sub>7</sub> are of about 1 to 3 carbons and include methyl, ethyl, iso-propyl and n-propyl.

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A particular group of compounds of formula (I) are those wherein X is oxygen and both R<sub>2</sub> and R<sub>3</sub>, and R<sub>4</sub> and R<sub>5</sub> together are methylenedioxy groups of the formula (II), wherein R<sub>6</sub> and R<sub>7</sub> are both hydrogen, both alkyl, or combine to form a spiro cyclopentyl or cyclohexyl ring, in particular where R<sub>6</sub> and R<sub>7</sub> are both alkyl such as methyl. A second group of compounds are those wherein X is CH<sub>2</sub> and R<sub>4</sub> and R<sub>5</sub> are joined to form a benzene ring. A third

group of compounds of formula (I) are those wherein both R2 and R3 are hydrogen.

The compounds of formula (I) may be synthesized by the following methods:

(a) Reaction of an alcohol of the formula RCH<sub>2</sub>OH with a chlorosulfamate of the formula CISO<sub>2</sub>NH<sub>2</sub> or CISO<sub>2</sub>NHR<sub>1</sub> in the presence of a base such as potassium a-butoxide or sodium hydride at a temperature of about -20° to 25° C and in a solvent such as toluene, THF or dimethylformamide wherein R is a moiety of the following formula (III):

$$R_5$$
 $R_4$ 
 $R_3$ 

15 (b) Reaction of an alcohol of the formula RCH<sub>2</sub>OH with sulfurylchloride of the formula SO<sub>2</sub>Cl<sub>2</sub> in the presence of a base such as triethylamine or pyridine at a temperature of about -40° to 25° C in a solvent such as diethyl ether or methylene chloride to produce a chlorosulfate of the formula RCH<sub>2</sub>OSO<sub>2</sub>Cl.

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The chlorosulfate of the formula RCH<sub>2</sub>OSO<sub>2</sub>Cl may then be reacted with an amine of the formula R<sub>1</sub>NH<sub>2</sub> at a temperature of abut 40° to 25° C in a solvent such as methylene chloride or acetonltrile to produce a compound of formula (I). The reaction conditions for (b) are also described by T. Tsuchiya et al. in Tet. Letters, No. 36, p. 3365 to 3368 (1978).

(c) Reaction of the chlorosulfate RCH<sub>2</sub>OSO<sub>2</sub>Cl with a metal azide such as sodium azide in a solvent such as methylene chloride or acetonitrile yields an azidosulfate of the formula RCH<sub>2</sub>OSO<sub>2</sub>N<sub>3</sub> as described by M.

Hedayatullah in Tet. Lett. p. 2455-2458 (1975). The azidosulfate is then reduced to a compound of formula (I) wherein R<sub>1</sub> is hydrogen by catalytic hydrogenation, e.g. with a noble metal and H<sub>2</sub> or by heating with copper metal in a solvent such as methanol.

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The starting materials of the formula RCH<sub>2</sub>OH may be obtained commercially or as known in the art. For example, starting materials of the formula RCH<sub>2</sub>OH wherein both R<sub>2</sub> and R<sub>3</sub>, and R<sub>4</sub> and R<sub>5</sub> are identical and are of the formula (II) may be obtained by the method of R. F. Brady in Carbohydrate Research, Vol. 14, p. 35 to 40 (1970) or by reaction of the trimethylsilyl enoi ether of a R<sub>6</sub>COR<sub>7</sub> ketone or aldehyde with fructose at a temperature of about 25° C, in a solvent such a halocarbon, e.g. methylene chloride in the presence of a protic acid such as hydrochloric acid or a Lewis Acid such as zinc chloride. The trimethylsilyl enol ether reaction is described by G. L. Larson et al in J. Org. Chem. Vol. 38, No. 22, p. 3935 (1973).

Further, carboxylic acids and aldehydes of the formulae RCOOH and RCHO may be reduced to compounds of the formula RCH2OH by standard reduction techniques, e.g. reaction with lithium aluminum hydride, sodium borohydride or borane-THF complex in an inert solvent such a diglyme, THF or toluene at a temperature of about 0° to 100° C, e.g. as described by H.O. House in "Modern Synthetic Reactions", 2nd Ed., pages 45 to 144 (1972).

The compounds of formula I: may also be made by the process disclosed US Patent: No.4,513,006, which is incorporated by reference herein.

The compounds of formula I include the various individual isomers as well as the racemates thereof, e.g., the various alpha and beta attachments, i.e., below and above the plane of the drawing, of R2, R3, R4 and R5 on the 6-membered ring. Preferably, the oxygens of the methylenedioxy group (II) are attached on the same side of the 6-membered ring.

The activity of the compounds of formula I in treating psonasis was first evidenced in clinical studies conducted to evaluate the efficacy of topiramate in treating epilepsy. At least three patients who coincidentally had psonasis reported that there was a marked reduction in the psonatic lesions. Therefore, preclinical in vitro studies were conducted to evaluate the effects of topiramate on keratinocyte function as a putative mechanism of action for its potential beneficial effects in treating psonasis. One of the hallmarks of psonatic lesions is hyperproliferative epidermal keratinocytes. In general, agents that affect the proliferation of keratinocytes have an inverse effect on differentiation, i.e. they would inhibit growth and enhance differentiation. Two measures of keratinocyte function were therefore evaluated: cell growth and differentiation.

In these studies, keratinocytes were grown in Medium154, a low calcium medium supplemented with bovine pituitary extract (BPE), bovine insulin, bovine transferrin, human epidermal growth factor (EGF) and hydrocortisone. Keratinocytes were grown to 60-80% confluence and subcultured by using trypsin/EDTA.

Four separate experiments were performed to evaluate the dosedependent effect of topiramate on keratinocyte cell growth as measured by maturity after six days of treatment. Topiramate was dissolved in DMSO to make a 100 mM stock solution. In all experiments the final concentration of DMSO in the cell incubation medium was 0.1%. Vehicle controls (0.1% DMSO) were included in each experiment. Cell growth was induced by a combination of the growth factors EGF and BPE. Topiramate had a modest inhibitory effect on cell growth under these assay conditions; however, no dose dependence was observed (R.W Johnson Pharmaceutical Research Institute Laboratory Notebook No. 12183 and 12540). The maximal response was observed at 10 micromolar, 32 ± 10% inhibition. While there was a trend toward inhibition of cell growth, this did not reach statistical significance (p>0.05).

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The effect of topiramate on keratinocyte differentiation was measured by the expression of transglutaminase-1 protein after three days of treatment. Three separate experiments were performed. Differentiation was evaluated in both low calcium and high calcium incubation conditions. An increase in differentiation would be most readily observed under low calcium conditions whereas an inhibition of differentiation could be detected under conditions of high calcium-induced differentiation. Topiramate caused a modest increase in transglutaminase-1 protein with both conditions, indicating an enhancing effect. A follow-up study was conducted which extended the incubation time to 5 days to look for further enhancement. No additional increases in transglutaminase-1 were observed in this latter study.

The results of these studies indicate that topiramate's effects on keratinocyte function are consistent with those expected for an agent that would affect the hyperproliferative keratinocyte response associated with psoriasis; inhibition of cell growth and enhancement of differentiation.

For treating psoriasis, a compound of formula (I) may be employed at a daily dosage in the range of about 50 to 400 mg administered orally, usually in two divided doses, for an average adult human. A unit dose would contain about 25 to 200 mg of the active ingredient. Alternatively, a compound of formula (I) may be administered topically to the affected area of the skin once or twice daily at a dosage in the range of 5 to 50 mg.

To prepare the pharmaceutical compositions of this invention, one or more sulfamate compounds of formula (I) are intimately admixed with a pharmaceutical carrier according to conventional pharmaceutical compounding techniques, which carrier may take a wide variety of forms depending on the form of preparation desired for administration, e.g., oral, by suppository, or parenteral. In preparing the compositions in oral dosage form, any of the usual pharmaceutical media may be employed. Thus, for liquid oral preparations, such as for example, suspensions, elixirs and solutions, suitable carriers and

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additives include water, glycols, oils, alcohols, flavoring ag nts, preservatives, coloring agents and the like; for solid oral preparations such as, for xample, powders, capsules and tablets, suitable carriers and additives include starches, sugars, diluents, granulating agents, lubricants, binders, disintegrating agents and the like. Because of their ease in administration, tablets and capsules represent the most advantageous oral dosage unit form, in which case solid pharmaceutical carriers are obviously employed. If desired, tablets may be sugar coated or enteric coated by standard techniques. Suppositories may be prepared, in which case cocoa butter could be used as the carrier. For parenterals, the carrier will usually comprise sterile water, though other ingredients, for example, for purposes such as aiding solubility or for preservation, may be included. Injectable suspensions may also be prepared in which case appropriate liquid carriers, suspending agents and the like may be employed.

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Topiramate is currently available for oral administration in round tablets containing 25 mg, 100 mg or 200 mg of active agent. The tablets contain the following inactive ingredients: lactose hydrous, pregelatinized starch, microcrystalline cellulose, sodium starch glycolate, magnesium stearate, purified water, carnauba wax, hydroxypropyl methylcellulose, titanium dioxide, polyethylene glycol, synthetic iron oxide, and polysorbate 80.

The pharmaceutical compositions herein will contain, per dosage unit, e.g., tablet, capsule, powder injection, teaspoonful, suppository and the like from about 25 to about 200 mg of the active ingredient.

1. A method for treating psoriasis comprising administering to a human afflicted with such condition a therapeutically effective amount for treating such condition of a compound of the formula I:

$$R_5$$
 $R_4$ 
 $R_2$ 
 $R_3$ 
 $R_2$ 

wherein

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X is CH2 or oxygen;

R<sub>1</sub> is hydrogen or alkyl; and

R<sub>2</sub>, R<sub>3</sub>, R<sub>4</sub> and R<sub>5</sub> are independently hydrogen or lower alkyl and, when X is CH<sub>2</sub>, R<sub>4</sub> and R<sub>5</sub> may be alkene groups joined to form a benzene ring and, when X is oxygen, R<sub>2</sub> and R<sub>3</sub> and/or R<sub>4</sub> and R<sub>5</sub> together may be a methylenedioxy group of the following formula (II):

20 wherein

R<sub>6</sub> and R<sub>7</sub> are the same or different and are hydrogen, lower alkyl or are alkyl and are joined to form a cyclopentyl or cyclohexyl ring.

- 2. The method of claim 1 wherein the compound of formula I is topiramate.
- 3. The method of claim 1, wherein the therapeutically effective amount is of from about 50 to 400 mg.
- 4. The method of claim 1, wherein the amount is of from about 25 to 200 mg.

# INTERNATIONAL SEARCH REPORT

International Application No. 97/10891

. CLASSIF	FICATION OF SUBJECT MATTER  1 K 31/35, A 61 K 31/36	
A 6.	1 K 31/35, A 01 K 31/30	
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C DOCUM	IENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to elaim No.
A	US 4513006 A	1-4
'	(MARYANOFF, B.E. et al.) 23 April 1985 (23.04.85),	a
	abstract, claims 5-9,	
•	1 column 1. lines $16-33$ .	
	column 3, line 15 - column 5,	
	line 19, example 3 (cited in the application).	
	(Cited in one appear	1,3,4
A	US 4792569 A	1,3,4
	(MARYANOFF, B.E. et al.) 20 December 1988 (20.12.88),	
ļ	abstract, claims 1,10-12,	1
	column 1, lines 20-37,	
	column 3, line 29 - column 4, line 33.	
	arther documents are listed in the continuation of box C. Patent family	members are listed in annex.
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### INTERNATIONAL SEARCH REPORT

Inter ional application No.

PCT/US 97/10891

Bxi	Observations where certain claims were f und unsearchable (Continuati n of item 1 f first sh et)
This Inte	emational Search Report has not been established in reepact of certain claims under Article 17(2)(a) for the following reasons:
1. X 2	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:  Remark: Although claim(s) 1-4  is(are) directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.  Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such
	an extent that no meaningful International Search can be carried out, specifically:
з. 🔲	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box ii	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inte	emational Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional eearch fees were timely paid by the applicant, this International Search Report covers all eearchable claims.
2.	As all eearchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional eearch fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Noe.:
Remark	on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

### ANHANG

zum internationalen Recherchen-bericht über die internationale Patentanmeidung Nr.

#### ANNEX

to the international Search Report to the International Patent Application No.

# ANNEXE

au rapport de recherche inter-national relatif à la demande de brevet internationai n°

### PCT/US 97/10891 SAE 168598

In diesem Anhang sind die Mitglieder der Patentfamilien der im obenge-nannten internationalen Recherchenbericht angeführten Patentdokumente angegeben. Diese Angaben dienen nur zur Unter-richtung und erfolgen ohne Gewähr.

This Annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The Office is in no way itable for these particulars which are given merely for the purpose of information.

La presente annexe indique les membres de la famille de brevets reiatifs aux documents de brevets cités dans le rapport de recherche international visée ci-dessus. Les reseignements fournis sont donnés à titre indicatif et n'engagent pas la responsibilité de l'Office.

In Recherchenbericht opeführtes Patentdokument Patent document cited in search report ocument de brevet cité ans ie rapport de recherche	Datum der Veröffentlichung Publication date Date de publication	Mitglied(er) der Patentfamilie Patent family member(s) Membre(s) de la familie de brevets	Datum der Veröffentilchung Publication date Date de publication	
S A 4513006	23-04-85	94213441111345411115145555404484253600404 48454889999000044442136699848525779888866447771212125555888666257700755658178202459 306447777888886664655555888866627700755658178202459 3552455999991111111155688 1052455999991111111155688 10524455999991111111155688 105524559999991111111155688 105524559999991111111155688 105524559999991111111155688 105524559999991111111155688 105524559999991111111155688 105524559999991111111155688 105524559999991111111155688 105524559999991111111155688 105524559999991111111155688 1055245599999911111111556888 1055245599999911111111556888 105524559999991111111556888 1055245599999991111111556888 10552455999999911111111556888 10552455999999999999999999999999999999999	85788451111CCCCCCCAA5995B557CCA744 868888899999998888888888888999998999898888	
US A 4792569	20-12-88	keine – none –	rien	